

proton channels from influenza A and B viruses by solution NMR spectroscopy. The channel structures reveal pore features that are important for proton gating and proton relay. Structural details of the anti-influenza drug, rimantadine, bound to the channel suggests an unexpected allosteric mechanism of drug inhibition and drug resistance, which has been verified by thorough functional and mutagenesis experiments.

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Proton Transport Through Channels: Insights and Surprises from Molecular Simulation

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The behavior of hydrated excess protons passing into and through transmembrane proton channels will be presented, based on the results of a novel multi-scale computer simulation methodology. The unique electrostatics related to the dynamic delocalization of the excess proton charge defect will be elaborated, as well as its effect on the channel proton transport and selectivity properties. The often opposing and asymptotic viewpoints related to electrostatics on one hand and Grotthuss proton shuttling on the other will be reconciled and unified into a single conceptual framework. Specific simulation results will be given for various channel systems, including the M2 channel of influenza A, proton selective mutant aquaporin-1 channels, the CIC Cl⁻/H⁺ antiporter, and models of the Hv1 voltage gated proton channel. Comparison to experimental results will be discussed where possible.

Platform L: Protein Folding Pathways

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Protein Folding: Independent Unrelated Pathways or Predetermined Pathway with Optional Errors

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There is a fundamental conflict between two different views of how proteins fold. Kinetic experiments and theoretical calculations are often interpreted in terms of different population fractions folding through different intermediates in independent unrelated pathways (IUP model). However, detailed structural information indicates that all of the protein population folds through a sequence of intermediates predetermined by the foldon substructure of the target protein and a sequential stabilization principle. These contrary views can be resolved by a predetermined pathway_optional error (PPOE) hypothesis. The hypothesis is that any pathway intermediate can incorporate a chance misfolding error that blocks folding and must be reversed for productive folding to continue. Different fractions of the protein population will then block at different steps, populate different intermediates, and fold at different rates, giving the appearance of multiple unrelated pathways. A test of the hypothesis matches the two models against kinetic folding results for two proteins, hen lysozyme and staphylococcal nuclease, which have been interpreted previously in terms of independent parallel pathways. Folding kinetics of both proteins fit equally well to the two models, indicating that the measured experimental data does not require alternative parallel pathways. The fitted PPOE reaction scheme leads to known folding behavior, whereas the IUP properties are contradicted by experiment. The appearance of a conflict with multipath theoretical models seems to be due to their different focus, namely on multitrack microscopic behavior versus cooperative macroscopic behavior. The integration of three well documented principles in the PPOE model (cooperative foldons, sequential stabilization, optional errors) provides a unifying explanation for how proteins fold and why they fold in that way.

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Multiple Routes and Milestones in the Folding of HIV-1 Protease Monomer

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Evolution has lead proteins to display funneled energy landscapes with small degrees of ruggedness. However, a funneled landscape does not preclude the

presence of multiple kinetically relevant folding routes. Here we show that for an extremely relevant biological case, the monomer of HIV type 1 protease (HIV-1-PR), multiple pathways and milestones can coexist along the folding process.

We provide a comprehensive picture of the folding mechanism of HIV-1-PR monomer using a variety of theoretical and computational techniques. These include all-atom molecular dynamics simulations in explicit solvent, an analysis of the network of structure clusters found in multiple high-temperature unfolding simulations and a complete characterization of the free energy surface carried out using an all-atom structure based potential and a combination of metadynamics and parallel tempering.

Our results confirm that the monomer in solution is stable and show unambiguously that at least two (un)folding pathways exist. Moreover, we demonstrate how the formation of a hydrophobic core can be considered a milestone in the folding process which must occur along all the routes that lead towards the protein's native state. These results also provide a theoretical framework that is able to rationalize both the experimental evidences and the evolutionary data for HIV-1-PR monomer. Finally, our characterization of the ensemble of possible folding routes substantiates a rational drug design strategy based on inhibiting the folding of each of the subunits that build the HIV-1 protease homo-dimer.

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Characterizing Energy Landscapes of Proteins and Identifying Shape-Determining Factors

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Protein folding is a long standing problem in biology, whose mechanism is still not completely understood. Funnel-shape energy landscape has been proposed as a plausible folding mechanism. However, the factors that determine the funnel-shape energy landscapes is largely unknown. In this study, we use hydrophobic-hydrophilic (HP) model to investigate the factors that affect the funnel-shape of protein energy landscapes. We designed a clustering method based on graph theory to analyze the conformations sampled using a recently developed Monte Carlo method, FRESS. We found that the way conformations move from one to another defined by a particular sampling method (move set) has a significant effect on the shape of protein energy landscapes. To our surprise, both protein-like sequences and random sequences with around 50% hydrophobic residues have a stable state represented by a single dominant cluster, consisting of a large number of similar conformations. The energy landscapes resemble a funnel, where there are many paths to minimum energy conformations in the dominant cluster from conformations of higher energies. We also found that sequences with hydrophobic residues above or below the optimal range around 50% do not have a single stable state. In stead, there are many much smaller clusters, representing multiple local energy minima. Our finding is consistent with the compositions of hydrophobic and polar residues in globular proteins (fold to unique structures) and intrinsically disordered proteins (IDPs). Our study suggests that in computational simulations, move sets affect significantly the shape of protein energy landscapes; hydrophobic interaction is likely a major force leading to the funnel-shape energy landscape of proteins; and the composition of hydrophobic and polar residues is an important sequence feature for the formation of funnel-shape of protein energy landscapes.

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Common Folding Mechanism of a Peptide Revealed by Multiple MD Simulations

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Here, we report on the folding of a 15 residue beta-hairpin peptide (Peptide 1) using multiple unbiased, atomistic molecular dynamics (MD) simulations. Fifteen independent MD trajectories, each 2.5 microseconds-long for a total of 37.5 microseconds are performed in explicit solvent, at room temperature and without the use of enhanced sampling techniques. The computed folding time of 1-1.5 microseconds obtained from the simulations is in good agreement with experiment. A common folding